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(54) A combination comprising candesartan and rosuvastatin for the treatment of atherosclerosis Zusammensetzung zur Behandlung von Atheroskierose, die Candesartan und Rosuvastatin enthält Combinaison comprenant du candesartan et de la rosuvastatine pour le traitement de l'athérosciérose

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(56) References cited: EP-A- 1 314 425 WO-A-01/76573 WO-A-02/058731 WO-A-2004/096810

 CHEN JIAWEI ET AL: "Marked upregulation of ilpoxygenase-1, a receptor for ox-low-density lipoprotein in atherosclerosis, and its total ablation by candesartan and rosuvastatin given concurrently: "JOURNALO PTHE AMERICAN COLLEGE OF CARBIOLOGY, vol. 43, no. 5 Supplement A, 3 March 2004 (2004-03-03), page 498A, XP002319611 & 53RD ANNUAL SCIENTIFIC ESSSION OF THE AMERICAN COLLEGE OF CARBIOLOGY, NEW ORLEANS, LA, USA; MARCH 07-10, 2004 ISSIN: 0735-097

 PATENT ABSTRACTS OF JAPAN vol. 2002, no. 09, 4 September 2002 (2002-09-04) & JP 2002 145770 A (SANKYO CO LTD), 22 May 2002 (2002-05-22)

Note: Within nine months of the publication of the mention of the grant of the European patent in the European Patent Bulletin, any person may give notice to the European Patent Office of opposition to that patent, in accordance with the implementing Regulations. Notice of opposition shall not be deemed to have been filed until the opposition fee has been paid. (Art. 99(1) European Patent Convention). [0001] The present invention relates to a combination comprising candesartan and rosuvastatin.

[0002] The present invention further relates to pharmaceutical compositions comprising the combination mentioned hereinbefore. The present invention further relates to the use of a combination mentioned hereinbefore in the manufacture of a medicament for the prevention or treatment of atherosclerosis.

[0003] Atheroscienosis is a condition mediated by complex pathological processes within result in irregularly distributed lipid deposits in the arteries and is a major contributory factor to coronary heart disease. A reduction in atheroscienosis is herefore a major target for reducing 15 the arteries are distributed in the contributed of cardiovascular events for example, myocardial infraction, worsening of angine, cardiac arrest, stroke, congestive heart failure and cardiovascular desth.

[0004] Dyslipidemia, particularly increased plasma 20 level of low-density lipoprotein (LDI) is one of the major risk factors in atherosclerosis. Clinical studies have demostrated that reducing plasma LDI level with 3-ty-drowy-3-methylipural yeenzyme A (HMG CoA) reductase inhibitors, commonly known as statins, results in a cover risk of aeriolyxeaulur eventive security and control security of the control security and control

[0005] Activation of the renin-englolensin system (RAS) may be considered another important risk factor in atheroscelerosis. Activation of RAS with the formation of anglotensin (II) (A (III) and the activation of A (III) recoptors have been implicated in atherogenesis, plaque unpture, myocardial ischemic dysfunction and congestive heart failure (Singh and Mehta, Arch Intern Med, 2003, vol 153, 1298-13904).

[0006] International Patent Application WO 96/26/188 of diadoses treatment of attenoscions with 4(II) resolved by diadoses vice and the foliation blockers, optionally in combination with HMGCoA reductase inhibitors. International Patent Application WO 17/95/37 discloses the use of a combination of at least two of an A(II) antagonist, an ACE (angiotensin conventing enzyme) inhibitor and an HMGCoA reductase with the prevention or delay of progression in a list of conditions, amonast which is attenoscherois.

[0007] We have surprisingly found that the combination of the A(II) antagonist candeseartan and the HMG
CoA reductase inhibitor resurseatath has a synergistic
effect in the reduction of atheroscienosis. This synergistic
effect appears to arise from synergistic inhibition of expression of a number of inflammatory mediators involved in the RAS (for example COA), metalloprotionisse (MMPs)) and/or inhibition of the expression of the receptor LOX-1 (which is a receptor for oxidised LDL. on
endothelial cells). The synergistic effect provides strong
evidence for cross-talk between the RAS and dyslipdemia in atheroscenesis.

[0008] It will be appreciated that the activity of MMPs may be regulated in-vivo by their tissue inhibitors (TIMPs). We have also shown that the expression of

TIMP-1 and TIMP-2 is up-regulated by high-cholesterol diet, and markedly attenuated by the combination of candesartan and rosavastatin. These data lend credence to the concept that the balance between MMPs and TIMPs is altered by high-cholesterol diet, and that this imbalance can be 'normalized' by the combination of an A(II) antaonist and a linici lowering a country.

[0009] In one aspect of the invention is provided a combination comprising candesartan, or a pharmaceutically receptable salt thereof, and rosuvastatin, or a pharmaceutically acceptable salt thereof, for use in the prevention or treatment of atherosclerosis.

[0010] In one aspect of the invention is provided a combination comprising candesartan, or a pharmaceutically acceptable salt thereof, and rosuvastatin, or a pharmaceutically acceptable salt thereof, for use in the prevention of cardiovascular events.

[0011] Such a combination may also be useful in the treatment or prevention of other diseases associated with these mediators, for example in inflammatory diseases or conditions, such as ischemia-reperfusion injury (to the heart, brain, kidneys, lungs and liver), radiation-induced injury, burn injury and peripheral vascular diseases,

[0012] Candesartan may suitably be in the form of canfeesartan, or in the pro-drug form candesartan cilexetil. These forms may be formulated with a further agent such as a diuretic such as hydrochlorothlazide (for example, as markdeted as Atacand PlusTM).

[0013] Where herein candesartan is referred to, this includes both candesartan and candesartan cilexetil.
[0014] Preferably the calcium salt of rosuvastatin,

[0014] Preferably the calcium salt of rosuvastatin, which may be referred to as rosuvastatin calcium, is used in the various aspects of the present invention.

[0015] In general, pharmaceutically-acceptable salts

include acid addition salts such as methanesulfonate, tosylate, a-glycerophosphate, furnarate, hydrochloride, citrate, maleate, tartrate and (less preferably) hydrobromide. Pharmaceutically-acceptable salts in general also include salts formed with phosphoric and sulfuric acid. Pharmaceutically-acceptable salts generally include base salts such as an alkali metal salt for example sodium, an alkaline earth metal salt for example calcium or magnesium, an organic amine salt for example triethylamine, morpholine, N-methylpiperidine, N-ethylpiperidine, procaine, dibenzylamine, N,N-dibenzylethylamine, tris-(2-hydroxyethyl)amine, tris(hydroxymethyl)methylammonium, N-methyl d-glucamine and amino acids such as lysine. There may be more than one cation or anion depending on the number of charged functions and the valency of the cations or anions.

[0016] Herein, where the term "combination" is used it is to be understood that this refers to simultaneous, separate or sequential administration. In one aspect of the invention "combination" refers to simultaneous administration. In an another aspect of the invention "combination" refers to separate administration. In a further aspect of the invention "combination" refers to sequential administration. Where the administration is sequential or s

is used.

arate, the delay in administering the second component should be such that both agents are present in the body so as to produce the synergistic effect of the combination.

[0017] In a further aspect of the invention is provided a pharmaceutical composition which comprises cande-sartan, or a pharmaceutically acceptable satit thereof, and rosuvastatin, or a pharmaceutically acceptable satit thereof, in association with a pharmaceutically acceptable sold the satit thereof, in association with a pharmaceutically acceptable of different or carrier for use in the prevention or treatment of atheroscierosis.

[0018] In a further aspect of the invention is provided a pharmaceutical composition which comprises candesartan, or a pharmaceutically acceptable satt thered, and rosuvastatin, or a pharmaceutically acceptable satt thereof, in association with a pharmaceutically acceptate 15 bid diluent or carrier for use in the prevention or reduction of risk of cardiovascular events.

[0019] The compositions described herein may be in a form suitable for oral administration, for example as a tablet or capsule, for parenteral injection (including intravenous, subcutaneous, intramuscular, intravascular or infusion) for example as a sterile solution, suspension or emulsion, for topical administration for example as an ointment or cream, for rectal administration for example as a suppository or the route of administration may be by direct injection into the tumour or by regional delivery or by local delivery. In other embodiments of the present invention the compounds of the combination treatment may be delivered endoscopically, intratracheally, intralesionally, percutaneously, intravenously, subcutaneous- 30 ly, intraperitoneally or intratumourally. In general the compositions described herein may be prepared in a conventional manner using conventional excipients or carriers that are well known in the art.

[0020] Sultable pharmaceutically-acceptable excipiare that carriers for a table fromulation include, for exampie, inert excipients such as lactose, sodium carbonate,
calcium phosphate or calcium carbonate, granulating
and dishtegrating agents such as com starch or alginic
acid; binding agents such as gelatin or starch; luchicating
agents such as gelatin or starch; luchicating
agents such as gelatin or starch; luchicating
agents such as a strip or propy 4-hydroxybenzoate, and anti-oxidatins, such as assorbic acid. Tabiet formulations may be uncosted or coated either to modfly their disintegration and the subsequent absorption of
the active ingredient within the gastrointestinal tract, or
to improve their stability andror appearance, in either
case using conventional coating agents and procedures
well known in the art.

[0021] Compositions for oral use may be in the form of hard gelatin capsules in which the active ingredient is mixed with an inset solid excipient, for example, calcium carbonate, calcium phosphate or kaosin, or as soft gelatin capsules in which the active ingredient is mixed with water or an oil such as peanut oil, liquid paraffin or olive oil. [0022] Candesarian is commercially availables as "At-acand" liva" in Sosuvastatin acticum is commercially availables as "Creative". Suitable formula-

tions for the present invention include those which are commercially available.

- [0023] Suitable dosages of each component of the combination are those of the marketed commercial products. Alternatively, the synergy between the components may allow a lower dosage of one or both components to be used. For example, a dose of 4mg, Bmg, 16mg, 32mg, or up to 160mg of candesartan in combination with a dose of 80mg, 40mg, 20mg, 10mg 5mg or 2.5mg of rosuvastatin may be used. It will be understood that any
- one of the doses of candesartan may be combined with any suitable dose of rosuvastatin.

 [0024] In one aspect, 80mg of rosuvastatin is used. In
- another aspect, 40mg of rosuvastatin is used. In a further aspect, 20mg of rosuvastatin is used. In a further aspect, 10mg of rosuvastatin is used. In a further aspect, 5mg of rosuvastatin is used. In a further aspect, 2.5mg of rosuvastatin is used. In a further aspect, 2.5mg of rosuvastatin is used.
- [0025] In one aspect, between 32 and 160mg, such as a about 64-96mg of candlesartan is used. Conventinely, about 12 mg of candlesartan is used. In another aspect, 32mg of candesartan is used. In a truther aspect, 16mg of candesartan is used. In a further aspect, 16mg of candesartan is used. In a further aspect, 16mg of candesartan is used. In a further aspect, 4mg of candesartan is used. In a further aspect, 4mg of candesartan is used.
- [0026] It will be appreciated that the pharmaceutical composition according to the present invention includes a composition according to the present invention includes a composition comprising candessartan or a pharmaceutically acceptable sait thereof and or systematically acceptable excipient or carrier. Such a composition, for example in a single or all formulation conveniently provides the therapeutic combination product of the invention for simultaneous administration in the prevention
- [0027] Preferably the two components of the combination are both administered orally.

or treatment of atherosclerosis.

- [0028] Preferably the two components of the combination are administered as a single oral formulation.

 [0029] Preferably the combination is formulated for
 - once-a-day dosing.

 [0030] Conveniently, the combination is formulated as a single tablet or capsule.
- 45 [0031] The dosages and schedules described herein-before may be varied according to the particular disease state and the overall condition of the paffect for exemple, it many be necessary or desirable to reduce the above-mentioned doses of the components of the combination for treatment in order to reduce to xichly. Dosages and schedules may also vary if, in addition to a combination treatment of the present invention, one or more additional chemotherapeutic agents are used. Scheduling can be dementioned by the practitioner who is treating any particular patient using his professional skill and it noveledge.
- [0032] A pharmaceutical composition according to the present invention also includes separate compositions comprising a first composition comprising candesartan

or a pharmaceutically acceptable salt thereof and a pharmaceutically-acceptable excipient or carrier, and a second composition comprising resuvastatin or a pharmaceutically acceptable salt thereof and a pharmaceutically-acceptable excipient or carrier. Such a composition 5 conveniently provides the therapeutic combination of the invention for sequential or separate administration in the synergistic prevention or treatment of atherosclerosis but the separate compositions may also be administered simultaneously.

[0033] In another aspect of the invention there is provided a combination comprising candesartan, or a pharmaceutically acceptable salt thereof, and rosuvastatin, or a pharmaceutically acceptable salt thereof, for use as a medicament for the prevention or treatment of atherosclerosis.

[0034] In another aspect of the invention there is provided a combination comprising candesartan, or a pharmaceutically acceptable salt thereof, and rosuvastatin. or a pharmaceutically acceptable salt thereof, for use as a medicament for the prevention of cardiovascular evente

[0035] In another aspect of the invention there is provided a combination comprising candesartan, or a pharmaceutically acceptable salt thereof, and rosuvastatin. or a pharmaceutically acceptable salt thereof, for use in the manufacture of a medicament.

[0036] In another aspect of the invention there is provided a combination comprising candesartan, or a pharmaceutically acceptable salt thereof, and rosuvastatin, 30 or a pharmaceutically acceptable salt thereof, for use in the manufacture of a medicament for the prevention or treatment of atherosclerosis.

[0037] In another aspect of the invention there is provided a combination comprising candesartan, or a phar- 35 Animal Model maceutically acceptable salt thereof, and rosuvastatin, or a pharmaceutically acceptable salt thereof, for use in the manufacture of a medicament for the prevention of cardiovascular events.

[0038] According to a further aspect of the present in- 40 vention there is provided a kit comprising a combination of candesartan or a pharmaceutically acceptable salt thereof, and rosuvastatin; or a pharmaceutically acceptable salt thereof, optionally with instructions for use in the prevention or treatment of atherosclerosis.

[0039] According to a further aspect of the present invention there is provided a kit comprising:

- a) candesartan in a first unit dosage form; b) rosuvastatin in a second unit dosage form; and
- c) container means for containing said first and second dosage forms; and optionally
- d) with instructions for use in the prevention or treatment of atheroscierosis

[0040] According to another aspect of the present invention there is provided a method of inhibiting expression of CD40, and/or metalloproteinases (MMPs) by administering a combination of candesartan, or a pharmaceutically acceptable salt thereof and rosuvastatin, or a pharmaceutically acceptable salt thereof.

[0041] Particular metalloproteinases are Mump-1. MMP-2 and MMP-9.

[0042] According to another aspect of the present invention there is provided a method of treating atherosclerotic patients by inhibition of expression of CD40 and/or metalloproteinases (MMPs) by administering an amount 10 of a combination of candesartan, or a pharmaceutically acceptable salt thereof and rosuvastatin, or a pharmaceutically acceptable salt thereof suitable for inhibition of expression of CD40 and/or metalloproteinases (MMPs).

[0043] Additionally, the invention can be useful in normalizing the balance between MMPs and TIMPS by administration of an amount of a combination of candesartan, or a pharmaceutically acceptable sait thereof and rosuvastatin, or a pharmaceutically acceptable salt

[0044] Additionally, the present invention can be useful in inhibiting expression of LOX-1 by administering a combination of candesartan, or a pharmaceutically acceptable salt thereof and rosuvastatin, or a pharmaceutically acceptable salt thereof.

[0045] Additionally, the present invention can be useful in treating atherosclerotic patients by inhibition of expression of LOX-1 by administering an amount of a combination of candesartan, or a pharmaceutically acceptable salt thereof and rosuvastatin, or a pharmaceutically acceptable salt thereof suitable for inhibition of expression of LOX-1.

Materials and Methods

[0046] Five pairs of C57BL/6J mice and three pairs of homozygous apo-E knockout mice (on C57BL/6J background) were obtained from Jackson Laboratories (Bar Harbor, ME). They were bred by brother-sister mating and housed in a room lit from 6:00 AM to 6:00 PM and kept at 21°C. The C57BL/6J mice (n=10) were continued on regular diet for the entire study period. The apo-E knockout mice were divided into four groups. Group 1 (n=10) animals were given high-cholesterol diet (1% cholesterol) alone for 12 weeks since the age of 6 weeks; Group 2 (n=10) animals were given high-cholesterol diet with candesartan (1mg/kg/d) for 12 weeks since the age of 6 weeks; Group 3 (n=10) animals were given highcholesterol diet with the rosuvastatin (1mg/kg/d) for 12 weeks since the age of 6 weeks; Group 4 (n=10) animals were given high-cholesterol diet with candesartan (1mg/kg/d) and rosuvastatin (1mg/kg/d) for 12 weeks since the age of 6 weeks.

[0047] At the end of 12-week-treatment, the mice were sacrificed and subject to studies described below. All experimental procedures were performed in accordance with protocols approved by the Institutional Animal Care and Usage Committee of University of Arkansas for Medical Sciences.

Quantitative Analysis of Atherosclerotic Plagues

[0048] At the end of 12-week-treatment, 5 mice from each group were euthanized and the aortas were separated from surrounding tissues. After removal of the adventifial fat issue, the aortas were opened longitudinally from the aorta and to the like biturcation, and tixed in 10% formalin for 24 hours. Then the aortas were rinsel in 70% alcohol fieldly, stained with Sudan IV Soution for 15 minutes, differentiated in 80% alcohol for 20 minutes and washed in nunning water for 1 hour (Russell L. Techniques for studying atheroselerotic lesion, Lab Invest. 1958; 742-47). The aortas were mounted and their pictures were taken with a camera connected to a dissection microscope. The images were analyzed by software (Image ne Pro Plus. Media Covbernelice).

RNA Preparation and Analysis BART-PER

[0049] At the end of 12-week-treatment, 5 mice from each group were euthanized and the aortas (from aorta arch to iliac bifurcation) were separated from surrounding 25 tissues and stored on dry ice. Each aorta was cut into four segments, two of which were used to extract total RNA with the single-step acid-quanidinium thiocyanatephenol-chloroform method as described earlier (27). One microgram of total RNA was reverse transcripted into cD- 30 NA with oligo-dT (Promega, Madison, WI, U.S.A.) and Maloney murine leukemia virus (M-MLV) reverse transcription (Promega) at 42°C for 1 hour. Two microliters of reverse transcription (RT) material was amplified with Tag DNA polymerase (Promega) and a primer pair spe- 35 cific to mouse LOX-1, CD40 or MUMPS (MMP-1, -2, -9). For mouse LOX-1, forward primer: 5'-TTACTCTCCAT-GGTGGTGCC-3', reverse primer: 5'-AGCTTCTTCT-GCTTGTTGCC-3' were used, 30 cycles of polymerase chain reaction (PCR) were performed at 94°C for 40 sec- 40 onds (denaturation), 55°C for 1 minute (annealing), and 72°C for 1 minute (extension). The size of polymerase chain reaction (PCR) product was 193 base pairs. For mouse CD40, forward primer 5'-GTTTAAAGTCCCG-GATGCGA-3' and reverse primer 5'-CTCAAGGCTAT- 45 GCTGTCTGT-3' were used. 35 cycles of polymerase chain reaction (PCR) were performed at 94°C for 1 minute (denaturation), 55°C for 1 minute (annealing), and 72°C for 1 minute (extension). The size of PCR product was 408 base pairs. For mouse MMP-1, forward primer 5'-GGACTCTCCCATTCTTAATGA T-3' and reverse primer 5'-CCTCTTTCTGGATAACATCATCA AC-3' were used. For mouse MMP-2, forward primer 5'-AT-CAAGGGGATCCAGGAGC-3' and reverse primer 5'-GCAGCGATGAAG ATGATAG-3' were used. For mouse 55 MMP-9, forward primer 5'-AGTTTGGTGTCGCGGAG-CAC-3' and reverse primer 5'-TACATGAGCGCTTCCG-GCAC-3' were used. For all MMPs, 35 cycles of PCR

were performed at 84°C for 1 minute (denaturation), 58°C for 1 minute (annealing), and 75°C for 1 minute (extension). The sizes of PCR product were 627,718 and 753 base pairs, respectively. A primer pair specific to mouse hackin was used as house-keeping gene (forward primer: 5°-TCTACAATGACTGCGTTG-3', reverse primer: 5°-TCTACAATGACTGCGTTG-3', acceptance used at 94°C for 30 seconds (denaturation), 55°C for 1 minute (annealing), and 72°C for 1 minute (extension). PCR product was 560 base pairs. The reverse transcription PCR (RT-PCR)-ampfiled sample was visualized on 1.5% agarose gu using ethildum bromide.

Protein Preparation and Analysis by Western Blot

[0050] Each mouse aorta was cut into four segments. Two of them were used to extract RNA, and the remaining two were used to extract protein as described previously (14). In brief, the aprilo tissues were homogenized and lysed in lysis buffer, then centrifuged at 4000 rpm for 10 minutes at 4°C. The lysate proteins from aortas (20 µg/ lane) were separated by 10% SDS-PAGE, and transferred to nitrocellulose membranes. After incubation in blocking solution (5% non-fat milk, Sigma), membranes were incubated with 1:750 dilution monoclonal antibody to mouse LOX-1, 1:500 dilution polyclonal antibody to mouse CD40 (Santa Cruz), 1µg/ml dilution monocional antibody to mouse MMP-1 (Oncogene), 1 µg/ml dilution monoclonal antibody to mouse MMP-2 (Oncogene). 1µg/ml dilution monoclonal antibody to mouse MMP-9 (Oncogene), 1:500 dilution polyclonal antibody to mouse TIMP- (Santa Cruz), 1:500 dilution polyclonal antibody to mouse TTMP-2 (Santa Cruz), or 1:5000 dilution monoclonal antibody to mouse β-actin (Sigma) for overnight at 4°C. Membranes were washed and then incubated with 1:5000 dilution specific secondary antibody (Amersham Life Science) for 2 hours at room temperature, and the membranes were washed and detected with the ECL system (Amersham Life Science). The relative intensities of protein bands were analyzed by Scan-gel-it software (Li DY, Zhang YC, Sawamura T, Mehta JL. Circ Res. 1999; 84:1043-1049).

Data Analysis

[0051] All data represent mean of duplicate samples. Data are presented as mean z SD. Statistical significance was determined in multiple comparisons among independent groups of data in which ANOVA and the F test indicated the presence of significant differences. A P value < 0.06 was considered significant.

Results

55 The synergistic anti-atherosclerotic effect of candesartan and rosuvastatin

[0052] Compared with the control mice (C57BL/6J

mice fed regular diet), the apo-E knockout mice fed highcholesterol diet developed extensive atherosclerosis (P<0.01 vs control mice). Although both candesartan and rosuvastatin alone decreased the extent of atherosclerosis (p<0.05 vs high-cholesterol diet alone), the combination reduced atherosclerosis to a much greater extent (P<0.05 vs candesartan or rosuvastatin alone plus highcholesterol del). Figure 1 shows results of representative experiments and the extent of atherosclerosis (mean ± 50) in different rorous of animals.

[0053] Candersartan and rosuvastatin alone decreased atherosclerosis by about 35% and 25% respectively. The combination reduced atherosclerosis by 70%, demonstrating a synergistic effect. This effect is illustrated graphically in Figure 2.

The synergistic effect of candesartan and rosuvastatin on LOX-1 expression

[0054] In the control C57BL/6J mice, the expression 20 of LOX-1 (mRNA and protein) was low. In contrast, LOX-1 expression (mRNA and protein) was markedly increased by high-choelasterol delt in spe-E knockout mice (P-0.01 ve control mice). Both candesartan and rosuvestatin alone decreased the LOX-1 expression (mRNA 25 and protein), albeit modestly (P-0.05 vs high-choissterol diet allone). The combination of candesartan and rosuvestatin had a dramatic inhibitory effect on the up-regulation of LOX-1 (mRNA and protein) in apo-E knockout mice (P-0.01 vs high-cholesterol diet allone).

The synergistic effect of candesartan and rosuvastatin on CD40 expression

[0055] Compared with the expression in control SCFBLC4 mice, CD40 expression (mRNA and protein) was markedly increased in apo-E knockout mice fed a high-cholesterol diet in (P-0.01 vs control mice). Although candesartan and rosuvastatin treatment alone slightly decreased CD40 expression (P-0.05 vs high-40-cholesterol diet alone), the combination of candesartan and rosuvastatin had a dramatic inhibitory effect on the up-regulation of CD40 (mRNA and protein) in the apo-E knockout mice (P-0.01 vs high-cholesterol diet alone).

The synergistic effect of candesartan and rosuvastatin on MMPs expression

[0056] Compared with the expression in control CSTBL/GJ mice, MMP-1, -2 and -9 expression (mRNA 50 and protein) was markedly increased in high-cholesterol diet-fed apo-E knockout mice (P-0.01 vs control mice). Both candesartan and rosuvastatin alone decreased MMP-1, -2 and -9 expression (mRNA and protein), albeit modestly (P-0.05 vs high-cholesterol diet alone). The combination of candesartan and rosuvastatin had a dramatic inhibitory effect on their expression (P-0.01 vs high-cholesterol diet alone).

The effect of candesartan and rosuvastatin on TRAPs expression

[0057] TIMP-1 and TIMP-2 protein expression was also increased in apo-E knockout mice by high-cholesterod diet (P-6.01 vs. control mice), but the increase was less than that of MMPs. Both candesartan and rosuvastatin alone reduced TEMP-1 and TIMP-2 expression by a small degree (P-0.05 vs. high-cholesterol diet alone), but the combination of candesartan and rosuvastatin had a greater inhibitor effect on their expression (P-0.01 vs.

high-cholesterol diet alone. P<0.05 vs. high-cholesterol

diet with candesartan or rosuvastatin).

Claims

- A combination comprising candesartan or a pharmaceutically acceptable salt thereof and rosuvastatin or, a pharmaceutically acceptable salt thereof for use in the prevention or treatment of atherosclerosis.
- A pharmaceutical composition which comprises a combination as claimed in Claim 1 in association with a pharmaceutically acceptable diluent or carrier for use in the prevention or treatment of atherosclerosis.
- Use of a combination as claimed in Claim 1 in the manufacture of a medicament for the prevention or treatment of atherosclerosis.
- A combination as claimed in Claim 1 wherein candesartan is in the form of candesartan cilexetil.
- [0055] Compared with the expression in control so 5. A combination as defined in Claim 1 for use in the prevention of cardiovascular events, such as myows markedly increased in apo-E knockout mice led a high-cholesterol diet in (P-0.01 vs control mice). At though candesartan and recovariating treatment alone under the compared of th
 - A pharmaceutical composition which comprises a combination as claimed in Claim 5, in association with a pharmaceutically acceptable diluent or carrier for use in the prevention of cardiovascular events.
 - Use of a combination as claimed in Claim 5 in the manufacture of a medicament for the prevention of carriovascular events
 - 8. A combination as claimed in Claim 5 wherein candesartan is in the form of candesartan cilexetil.
 - 9. A combination as claimed in any one of claims 1, 4, 5, and 8 wherein the candesartan is formulated with hydrochlorothiazide.

 7. A combination as claimed in any one of claims 1, 4, 5, and 8 wherein the candesartan is formulated with hydrochlorothiazide.

Patentansprüche

 Kombination, enthaltend Candesartan oder ein pharmazeutisch annehmbares Salz davon und Rosuvastatin oder ein pharmazeutisch annehmbares
 Salz davon zur Verwendung bei der Prävention oder Behandlung von Atherosklerose.

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- Pharmazeutische Zusammensetzung, enthaltend eine Kombination nach Anspruch 1 zusammen mit 10 einem pharmazeutisch annehmbaren Verdünnungsmittel oder Träger zur Verwendung bei der Prävention oder Behandlung von Atherosklerose.
- Verwendung einer Kombination nach Anspruch 1 bei 15 der Herstellung eines Medikaments zur Prävention oder Behandlung von Atheroskierose.
- Kombination nach Anspruch 1, wobel das Candesartan in Form von Candesartancilexetil vorliegt.
- Kombination nach Anspruch 1 zur Verwendung bei der Prävention von Herzkreislaufereignissen wie Herzinfarkt, einer Verschlimmerung von Angina, Herzstillstand, Schlaganfall, dekompensierter Herzinsuffizienz und Herzkreislauftod.
- Pharmazeutische Zusammensetzung, die eine Kombination nach Anspruch 5 zusammen mit einem pharmazeutisch annehmbaren Verdünnungsmittel ³⁰ oder Träger enthält, zur Verwendung bei der Prävention von Herzkreislaufereignissen.
- Verwendung einer Kombination nach Anspruch 5 bei der Herstellung eines Medikaments zur Pr\u00e4vention 35 von Herzkreislauferelgnissen.
- Kombination nach Anspruch 5, wobei das Candesartan in Form von Candesartancilexetil vorliegt.
- Kombination nach einem der Ansprüche 1, 4, 5 oder 8, wobei Candesartan mit Hydrochlorothiazid formuliert ist.

Revendications

- Combinaison comprenant le candésartan ou un sel pharmaceutiquement acceptable de celui-cì et la rosuvastatine ou un sel pharmaceutiquement acceptable de celle-ci, destinée à être utilisée dans la prévention ou le traitement de l'athéroscéirose.
- Composition pharmaceutique, caractérisée en ce qu'elle comprend une combinaison selon la revendication 1 en association avec un diluant ou un support pharmaceutiquement acceptable, destinée à être utilisée dans la prévention ou le traitement de

l'athérosclérose

- Utilisation d'une combinaison selon la revendication 1, dans la fabrication d'un médicament destiné à la prévention ou le traitement de l'athérosclérose.
- Combinaison selon la revendication 1, caractérisée en ce que le candésartan est sous forme de candésartan cilexétil.
- Combinaison telle que définie dans la revendication 1, destinée à être utilisée dans la prévention d'événements cardiovasculaires tels que l'infarctus du myocarde, l'aggravation de l'angine, l'arrêt cardiaque, l'accident vasculaire cérébral, l'insulfisance cardiaque congestive et la mot cardiovasculaire.
- 6. Composition pharmaceutique, caractérisée en ce qu'elle comprend une combinaison selon la revendication 6, en association avec un diluant ou un support pharmaceutiquement acceptable, destinée à être utilisée dans la prévention d'événements cardiovaculaires.
- Herzstillstand, Schlaganfall, dekompensierter Herz
 25 7. Utilisation d'une combinaison selon la revendication finsuffizienz und Herzkreislauftod.

 5, dans la fabhication d'un médicament destiné à la prévention d'événements cardiovasculaires.
 - Combinaison selon la revendication 5, caractérisée en ce que le candésartan est sous forme de candésartan cllexétil.
 - Combinaison selon l'une quelconque des revendications 1, 4, 5 et 8, caractérisée en ce que le candésartan est formulé avec de l'hydrochlorothiazide.

7

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Figure 1

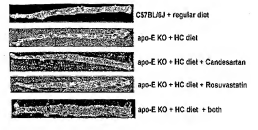
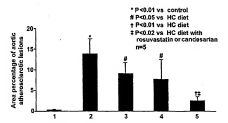


Figure 2



- 1. Control mice fed with regular diet
- 2. Apo-E KO mice fed with HC diet
- 3. Apo-E KO mice fed with HC diet together with rosuvastatin
- 4. Apo-E KO mice fed with HC diet together with candesartan
- 5. Apo-E KO mice fed with HC diet together with rosuvastatin and candesartan

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REFERENCES CITED IN THE DESCRIPTION

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